AMPLIFYING EXPRESSED SEQUENCES FROM GENOMIC DNA OF HIGHER-ORDER EUKARYOTIC ORGANISMS FOR DNA ARRAYS

FIELD OF THE INVENTION

[0001] The present invention relates to a method and devices that embody the method for in vitro amplification of expressed sequences directly from genomic DNA (gDNA) of all mammalian and/or higher-order plant species for DNA array fabrication. The method can be used to selectively amplify nucleic acid sequences, which contain sequence variations such as point mutations, deletions and insertions.

BACKGROUND

[0002] High-density arrays (HDAs) of cDNA or oligonucleotide have been powerful tools for profiling gene expression of particular cell or tissue types. Researchers have employed HDAs in their studies to uncover relationships between known genes, as well as, to reveal the function of previously uncharacterized genes. In current HDAs, the expressed genetic sequences, which are printed on the solid surfaces that form the arrays, typically come in two basic forms, selected from either 1) DNA fragments amplified from cDNA clones or genomic DNA of single cell organisms, or 2) synthetic oligonucleotides.

[0003] The current technology, while useful, has many associated problems, in particular regarding the amplification of cDNA fragments from cDNA clones. First is the issue of availability. Good cDNA samples are more cumbersome to procure. In cDNA samples procured commercially, about 30 percent of the clones contain

inaccurate or wrong identities, which makes them not useful and difficult, if not impossible, to amplify by polymerase chain reaction (PCR). Hence, one is forced to order multiple clones for a single gene. This is not cost effective and can lead to experimental errors. Further, many genetic clones are not available commercially. It is estimated that expressed-sequence tag (EST) clones represent less than about 80% mammalian genes. Second, the entire sequence for clones having inserts that are longer than about 500 base pairs (bp) in size is often unknown. It is likely that some chimeric and/or large-intron-containing fragments may be introduced into these sequences. This is problematic, since one segment may contain sequences from two different genes, which could result in misleading data and lead to wrong interpretations. The resulting difference in size between individual cDNA fragments could be over 5-fold. This amount of deviation can produce unacceptable degrees of variation in the experimental data. Third, a high level of background signal can result since all EST sequences contain poly-adenine (poly-A), which can bring about increased levels of false hybridization and is detrimental for detection.

[0004] An alternative approach to amplified cDNA fragments uses reverse transcription (RT) products of messenger RNA (mRNA) as templates for polymerase chain reaction (PCR), i.e., RT-PCR. The problem with this approach, however, is that only about 10% to 20% of genes are expressed in a given cell or tissue type. To amplify cDNA fragments for all genes, a comprehensive collection of mRNAs from various cells or tissues and different stages of development is a must. This kind of comprehensive collection is very difficulty to obtain given current technology. In addition, this approach is severely limited in its potential to study unclonable sequences. Hence, a need exists for a new method that can amplify all kinds of gene sequences, both known and hypothetical.

SUMMARY OF THE INVENTION

[0005] The present invention addresses the need for a simpler, yet more efficient method of amplifying gene sequences in mammalian and/or higher-order plant species. The method provides a means for large-scale production of genomic DNA (gDNA) sequences. The method comprises several steps. First, a 3'UTR of a gDNA sequence based on the presence of a stop codon and a polyadenylation signal in the gDNA

sequence corresponding to an expressed mRNA sequence is identified. Alternatively, a "hypothetical" whole or partial exon from a gene defined by computer software can also be used. A predetermined gDNA sequence within the 3'UTR is then selected, preferably using computer software. The predetermined gDNA sequence has an overall homology of less than or equal to about 40% to any other genomic sequence in the same genome. A probe for the predetermined gDNA sequence is designed. Next, a first polymerase chain reaction (PCR) of the 3'UTR on gDNA to generate PCR-product is performed, followed by segregating the resultant PCR-product by a size-separation process selected from the group consisting of electrophoresis and chromatography. The predetermined gDNA sequence within the 3'UTR has a length of about 200 to about 600 nucleotide bases. A predetermined band from the size-differentiated samples is chosen, and a second polymerase chain reaction is performed to amplify the sample. The method can generate large quantities of gDNA probes, which enables greater efficiency for printing in microarray formats.

[0006] The present invention also includes a biological array. The biological array comprises a substrate and deposited on the substrate a set of amplified gDNA fragment sequences generated according to the method above. Each amplified sequence is derived from the sequence of at least one exon, or a partial exon, and contains no polyadenosine nor requires a vector sequence.

[0007] Additional features and advantages of the present invention will be disclosed in the detail description that follows.

BRIEF DESCRIPTION OF THE DRAWINGS

[0008] FIGURE 1 is a schematic that illustrates the 3'UTR of a gene defined by the presence of a translational stop codon and polyadenylation (polyA) signal, as well as its relative location on the human genome. The boxes, on the left, represent exons. The longer open box, on the right, represents the last exon containing the 3'UTR. GSP stands for gene specific primer.

[0009] FIGURE 2 is a flowchart to demonstrate how to define a unique sequence within the 3'UTR of a gene and design a pair of primers for PCR amplification of the sequence directly from genomic DNA.

[0010] FIGURE 3 is a schematic representation of a flowchart for PCR amplifications. The basic steps are listed along the center. The schematic at left shows the strategy using T7/T3 primers for the second PCR, while the schematic at right shows the strategy using gene specific primers (GSP) for both rounds of PCR.

[0011] FIGURE 4 shows size distribution of the 3'UTR for 117 genes. Genes are classified along the X-axis into three groups based on the size of their 3'UTR: 1) < 200 bp, 2) between 200 to 400 bp, 3) > 400 bp. The number within each bar represents the number of genes within each group (Y-axis).

[0012] FIGURE 5A is an image of an agarose gel of the PCR products from the first round for 12 genes. The number of each sample is indicated along the top, and flanked on each side by a molecular weight marker graded in increments of 100 bp (ladder). The 600 bp band is indicated by a line with an arrow head.

[0013] FIGURE 5B is another image of an agarose gel of the PCR products from the second round for 24 genes. The number of each sample is indicated along the top. The molecular weight marker in increments of 100 bp (ladder) is shown at right. A line with an arrowhead indicates the 600-bp band.

DETAILED DESCRIPTION OF THE INVENTION

Definitions

[0014] The term "alternatively spliced messages," as used in the context of the present invention, refers to mature mRNAs originating from a single gene with variations in the number and/or identity of exons, introns, and/or intron-exon junctions.

[0015] The term "biosite" as used herein means a discrete area, spot or site on the active surface of an array, or base material, comprising at least one kind of immobilized biological material for use as a probe or other functionality.

[0016] The term "chimeric," as used in the context of the present invention, describes genes or constructs wherein at least two of the elements of the gene or construct – such as a sequence from one gene linked or physically connected with a sequence from another gene – are heterologous to each other.

[0017] The term "gene," as used in the context of the present invention, encompasses all regulatory and coding sequences contiguously associated with a single hereditary unit with a genetic function. Genes comprise exons (coding sequences) that may be

interrupted by introns (non-coding sequences). Genes can include non-coding sequences that modulate the genetic function, which includes, but is not limited to, those that specify polyadenylation, transcription regulation, DNA conformation, chromatin conformation, extent and position of base methylation and binding sites of proteins that control all of these. A gene's genetic function may require only RNA expression or protein production, or may only require binding of proteins and/or nucleic acids without associated expression.

[0018] The term "gene family," as used in the context of the present invention, refers to a group of functionally related genes, each of which encodes a separate protein.

[0019] The term "heterologous sequence," as used herein, refers to genetic sequences that are not operatively linked, or in nature are not contiguous to each other.

[0020] The term "homologous gene" or "homologous sequence," as used herein, refers to a gene that shares sequence similarity with the gene of interest. This similarity may be only a fragment of the sequence and often represents a functional domain, such as a DNA binding domain, a domain with tyrosine kinase activity, or the like. The functional activities of homologous genes are not necessarily the same.

[0021] The term "public sequence," as used herein, refers to any sequence that has been deposited in a publicly accessible database. This term encompasses both amino acid and nucleotide sequences. Such sequences are publicly accessible on the websites of the National Center for Biotechnology Information (NCBI), for example in the UniGene database (http://www.ncbi.nlm.nih.gov/UniGene). The UniGene database uses accession numbers assigned by NCBI as a unique identifier for each sequence in the databases, thereby providing a non-redundant database for sequences from various databases, including GenBank, EMBL, DBBJ (DNA Database of Japan), PDB (Brookhaven Protein Data Bank) and other like databases. The Basic Local Alignment Search Tool (BLAST) database (http://www.ncbi.nlm.nih.gov/BLAST) is used for searching.

[0022] The term "regulatory sequence," as used herein, refers to any nucleotide sequence that influences transcription or translation of initiation and rate, and stability and/or mobility of the transcript or polypeptide product. Regulatory sequences include, but are not limited to, promoters, promoter control elements, protein binding sequences,

5' and 3' UTRs, transcription start site, termination sequence, certain sequences within a coding sequence, polyadenylation sequence, introns, etc.

[0023] The term "related sequences," as used herein, refers to a nucleotide sequence that exhibits some degree of sequence similarity with another sequence.

[0024] The term "sequence tagged site" (STS), as used herein, refers to a short DNA sequence that has a single occurrence in the human genome and whose location and base sequence is known. Detectable by polymerase chain reaction (PCR), STSs are useful for localizing and orienting the mapping and sequence data that are reported from many different laboratories and serve as landmarks on the developing a physical map of the human genome. Many STSs are derived from bacterial artificial chromosome (BAC) and/or P1 (bacterial phage) artificial chromosome (PAC) end sequences. Expressed sequence tags (ESTs) are STSs derived from cDNAs.

[0025] The term "untranslated region" (UTR) is a contiguous series of nucleotide bases that is transcribed, but not translated during synthesis of a peptide or protein. These untranslated regions may be associated with particular functions such as increasing mRNA message stability. Examples of UTRs include, but are not limited to polyadenylation signals, termination sequences, sequences located between the transcription start site and the first exon (5'UTR) and sequences located between the last exon and the end of the mRNA (3'UTR), including regulatory sequences.

Description

[0026] The method and devices embodying the method of the present invention circumvents the problems associated with generating cDNA fragments from DNA clones or long oligonucleotides. The present method enables one to perform large-scale amplification of expressed sequences directly from mammalian genomic DNA (gDNA) as the starting material. This feature is an advantage, since gDNA is easier to obtain than RNA for more genetic sequences. The present method generally abstains from using clonal DNA (cDNA) or RNA-derived sequences. Rather, by means of simple PCR amplifications without cloning, the method produces amplified sequences that have greater specificity and size consistency than that observed with cDNA fragments, and allows for greater signal sensitivity than oligonucleotides.

[0027] PCR amplification of expressed sequences from gDNA of prokaryotic organisms, such as bacteria, and lower-order eukaryotic organisms, such as yeast, has been a relatively simple task. This is because, at about 100-1000 times smaller than the genome of humans or other mammalian species, the genome of prokaryotes and lowerorder eukaryotes are relatively simple and do not have repetitive sequences or virtually no introns. (Yeast has only three genes that are found to contain small introns.) To do PCR amplification directly from gDNA of mammalian or other higher-order eukaryotes has been traditionally either nearly impossible or fraught with great difficulties. In contrast to single cell organisms, mammalian or higher-order eukaryote genomes are much more complex, possessing many intron segments that divide gene sequences into multiple exons and many more, longer regulatory sequences. During the natural transcription and gene expression process, a precursor RNA containing both exons and introns is first transcribed. The introns are removed subsequently through splicing to form mRNA, i.e., expressed sequences. The presence of multiple introns often complicates the task for researchers to amplify coherent, accurate, expressed gene sequences by means of PCR amplification.

[0028] With PCR, it is possible to amplify a single copy of a specific target sequence in gDNA to a level detectable by several different methodologies. For instance, the methods may include hybridization with a labeled probe; incorporation of biotinylated primers followed by avidin-enzyme conjugated detection; or, incorporation of ³²P labeled deoxynucleotide triphosphates into the amplified segment. Although PCR amplification of human genomic DNA has been used to identify sequence-tagged sites (STS), simple sequence length polymorphism (SSLP), single-stranded sequence conformation polymorphism (SSCP), or single nucleotide polymorphism (SNP) when the sequence for the region of interest is available, the applications that use these kinds sequences do not need large quantities of the PCR products, as would be required in the preparation of DNA microarrays. Indeed, even though some have suggested using amplified human gDNA with primer pairs to generate STS probes, whereby selected primer pairs corresponding to the 3'UTR of gene transcripts are employed, it is doubtful that they can generate sufficient amounts of amplified product. This is so because of two basic factors. One, primers adapted from STS do not have the specificity designed for gDNA amplification, which can not effectively control for the

guanine-cytosine (G-C) content or overall quality of the primers. Two, a direct use of STS from gDNA for PCR reactions raises the potential for contamination by the gDNA in the preparations, which can lead to greater background or mismatched-hybridization signal. Furthermore, a detailed methodology is lacking.

[0029] More importantly, the applications that use the kinds of sequences discussed

tend to be indiscriminate about which particular sequence or region of gDNA is used; that is, these applications do not necessarily select for expressed gDNA sequences. which is a particular subpart of coding regions in a gene. Rather, expressed and nonexpressed sequences alike may be mixed together with no particular specificity. For the purposes of the present invention, to amplify expressed sequences from genomic DNA is usually difficult without previous knowledge of the intron/exon boundaries for a given gene. Mammalian introns often range in size from less than about 100 to over 10,000 base pairs (bp). The distance between two exons could be too long to be amplified by a regular PCR, and one or both primers could cross the boundary of two exons. This characteristic makes it very difficult for PCR process to work. [0030] Although no systematic study has been conducted on the genomic structure of the 3'untranslated region (3'UTR) for all known genes, numerous studies of the genomic structure for various genes indicate that the 3'UTR often exists as a single exon. Typically, the 3'UTR is the longest exon and forms part of all expressed sequences in gDNA. The 3'UTR is very specific, containing within it a unique sequence for each given gene. This phenomenon makes the 3'UTR a valuable tool to differentiate individual genes within a gene family. While not intending to be bound by theory, it is believed that one can amplify the 3'UTR from genomic DNA without having to rely on any information regarding the intron/exon boundaries. The 3'UTR can unlock the potential for high-throughput amplification of DNA sequences directly from gDNA, for the purpose of using gDNA in high volumes in the fabrication of highdensity microarray products according to the present invention.

[0031] The method of the present invention, having been developed according to the principle described above, has the following protocol. First, a gene having a known public sequence is derived from a publicly accessible database, such as the UniGene database, and analyzed using a pair wise search by means of BLAST. A 3'UTR or an exon of that gene is defined or identified by the length between the translational stop

codon (e.g., TAA, TGA, or TAG) and the last nucleotide before a polyadenylation signal (e.g., AATAAA or ATTAAA). For the present method to work more effectively, the 3'UTR should have a length of about at least 200 nucleotide bases. Second, a segment of sequence within the 3'UTR, ranging from about 75 to about 2000 nucleotide bases is further selected by BLAST-searching the original gene sequence against the entire UniGene database using a gene- or oligo-designer computer software program. Selected sequences have preferably about 200 nucleotide bases or less, to about 800 nucleotide bases or more. More preferably, the selected sequence has a length of about 200 bases to about 500 or 600 bases, more preferably from about 225 or 250 bases to about 400 or 450 bases. The purpose of this second step is to minimize homologous sequence that may be otherwise also selected for in the PCR process. Thus, the accuracy and efficiency of downstream PCR amplification is improved. Generally the less homologous, or more heterologous, the sequence is to other sections of the genome the better to reduce mismatches during hybridization. The homology of the segments as used herein is determined on an overall scale comparing the selected gene sequence to all other gene sequences of the genome. That is, no clustering occurs preferably in any one region, but is rather diffused throughout the sequence. The selected gDNA segment has an overall amount of homology of less than or equal to about 70% for highly homologous gene families, but is more commonly less than or equal to about 40%. Preferably, the overall homology is about 35% to about 20%-15% or less. Use of gene-designer computer software also permits one to pick the PCR segments in a high throughput mode, so that one can select segments of sequences for PCR in a large-scale and automated fashion. Figure 1 illustrates the process described above in schematic form, and Figure 2 further describes the process in a flow chart. [0032] Third, a primer design software, like web-based Primer 3 (http://www.genome.wi.mit.edu/cgi-bin/primer/primer3 www.cgi), is used to design a complement for the selected or predetermined gDNA sequence. The primers in reaction, in contrast to STS probes that are spotted on a surface, are designed with greater specificity for gDNA amplification according to more stringent parameters in terms of sequence length and about 50-60% G+C content. Individual primers are verified by BLAST search for correct gene origin and absence of random overlapping sequences. Generally, the primer designed for a given segment should not contain a

related sequence. Table 2 lists all primer sequences used. Two types of primer pair were designed at about 500 bp apart (or within 200-400 bp when the 3'UTR is less than 500 bp long) and away from repetitive sequences. Type I contains a T7 promoter at the 5'end of the gene specific primer (GSP) in the sense direction and a T3 promoter at the 5'end of the GSP in the anti-sense direction. In particular, the sequence for T7 promoter is 5'-TAATACGACTCACTATAGGG-3' and for T3 promoter is 5'-ATTAACCCTCACTAAAGGGA-3' (derived from InvitrogenTM). Type II primers only contain gene specific sequences. All primers were purchased from Sigma-GenosysTM as desalted and dried pellets. Each pellet was dissolved in ddH₂O to a final concentration of 500 μM.

[0033] Next, a first round of PCR is preformed under predetermined conditions, which will be explained more fully in the Experiments section, below. Two different strategies were applied. As shown in Figure 3, the flowchart, Strategy 1 is to employ Type I primers (GSP with T7 or T3 promoter at a 5'end) for the first PCR, then use T7 and T3 primers for the second PCR (Figure 3, left panel). The other, Strategy 2, is to use the same pair of gene specific primers, Type II primers (GSP alone), for both first and second round of PCR (Figure 3, right panel).

[0034] Generally, the PCR product from this first round are then separated according to size-differentiation. Various size-differentiation processes, such as electrophoresis or chromatography (e.g., High Performance Liquid Chromatography), may be used. The size-differentiated sequence sample or band of interest is then gathered up by a transfer pipette, without need for purification – this is, without the need to remove each sequence-band from its gel bed – and suspended in a small volume (~50 μL) of water. [0035] A second round of PCR is performed on the predetermined sequence sample under the same conditions as in the first round of PCR. The PCR product from this second round is subjected to column purification or gel electrophoresis to clean up the amplified sequences using a commercial purification kit and eluted into a final volume. [0036] The final amplified sequence(s) derived according to the method can be printed or otherwise deposited as an array of biosites on a treated glass (e.g., borosilicate, aluminosilicate, fused silica, treated with a propylsilane or the like), polymer (e.g., polystyrene or polypropylene, nylon filter), or metallic (e.g., gold, platinum, chromium, or silicon) substrate for DNA micro-assay purposes. These kinds of arrays are the

functional heart of DNA microarrays used in genomic studies, drug discovery, and other biological assays. The device can be characterized as having a set of gDNA fragments having the sequence of one exon having no poly-adenosine nor vector sequence, and having a sequence length that range from about at least 75-80 bases to about 1800-2000 bases. Preferred fragment lengths are about 200 to about 600 or 800 nucleotides. Particular uses and means of fabrication of specific arrays are described in detail in International Patent Application No. WO 00/77257, entitled "Gene Specific Arrays and the Use Thereof," by Narayan Baidya *et al.*, the complete contents of which are incorporated by reference into the present disclosure.

[0037] The DNA fragments, generated according to the present invention, function essentially like cDNA fragments that have been amplified from cDNA clones, but provide many advantages with few of the associated drawbacks. The present invention solves the procurement problem, since the method is not limited by or dependent on the availability of cDNA clones, nor does it depend on bacterial cultures. Hence, with gDNA fragments generated according to the present invention, it is possible to cover the entire mammalian genome. The method has an overall shorter processing time than current methods since it requires neither cloning nor initial purification after the first round of PCR. Using the method, one can maintain quality control relatively easily. Partially as a result of prior determination and size-differentiation, the final expressed gDNA sequences generated and amplified according to the inventive method have small size variations between individual amplified strands and no poly-adenosine sequences. This feature promotes more functional consistency in the amplified sequences. Further, in operation, they do not require vector sequences.

[0038] The method described here can be used widely to amplify expressed sequences from the genomic DNA of humans and other mammalian animals, as well as higher order plants. With the recent completion of sequencing of the entire human genome and of many other mammalian genomes, the intron/exon boundaries for all genes will soon be known. Since there is always one or multiple exons with a size longer than about 500 bp, the length of the 3'UTR will no longer be a limiting factor. All expressed sequences for virtually all genes can be amplified using this method. Even genes with currently hypothetical exons can be identified through use of the present invention. The sequence for hypothetical exons can be defined by computer software.

Even though predicted by gene prediction software, many genes in these genomes, however, may not be clonable – thus, not available as cDNA clones. At present, the only way to study unclonable sequences of genes is to use synthetic oligonucleotides. The present method amplifies expressed sequences of gDNA of at minimum about 75 bases – preferably about 200 bp – or longer, providing better performance than oligonucleotides, which can not provide sufficient signal due to their limited lengths (< 100-150 bp).

[0039] When amplifying expressed sequences from genomic DNA, a major issue is how to procure a sufficient amount of PCR fragments to print arrays on surfaces. The PCR amplification process is known to reach a plateau concentration of specific sequences. The human genome has about 3.2 billion base pairs. The amount of unique 1000 bp sequence within 10 μg of total genomic DNA is estimated to be about 0.32 pg. A single run of a single PCR reaction under a standard condition, i.e., to use 1 µg genomic DNA in 50 µL reaction for 35 cycles, usually yields less than 1 µg of PCR product at most. Multiple reactions will consume great amounts of gDNA, which is quite expensive. Hence, a second round of PCR is usually necessary to secure a sufficiently large quantity. Performing a second round of PCR using the first PCR product as templates directly, without purification, however, traditionally results in high background, which are seen as a big smear around the specific PCR product, as mentioned above. This phenomenon suggests that the presence of irrelevant sequences that may cause researchers to misinterpret the data from subsequent array analysis. [0040] The present method alleviates these problems – minimizing, if not eliminating them – through several advantageous features. It is believed that due partially to sizedifferentiation and one or more second round(s) of PCR, the present invention can produce at least about twice – if not three to five times or more – the amount of amplified product than that which can be attained through use of other ways of generating probes. The strands of amplified sequences generated using the present method are relatively size constant. Moreover, because gDNA does not contain polyadenosine sequences, nor undergoes polyadenylation, which is a posttranscriptional process, there is little likelihood of false hybridization. Since there is no poly-A to remove, the method saves time in the process.

[0041] The most commonly used protocol, currently available, to generate large amounts of gene specific PCR products is to perform a so-called nested PCR. That is, perform a first round of PCR with a pair of GSPs, and then a second round of PCR using another pair of internal GSPs. According to this procedure, each gene needs four GSPs for the PCR. The protocol, thus, creates more work in the design of the primer and also doubles the cost. This means that researchers need to design two pairs of primers, which is a possible limitation to the process. It is difficult to find a second pair of primers within the segment defined by the first primer pair.

[0042] An approach, practiced in small scale laboratory work, is to perform a first round of PCR, cut-out a gel slice containing the products from the first PCR, purify the DNA using commercially available kits, and then use it as templates for the second round of PCR. This process, however, is time consuming. The inventive method eliminates the need for a purification step, which is one of its important improvements over the prior art, and enables large-scale production of large amounts of amplified sequence in a high-through-put manner for DNA microarrays. Instead of using a laser bladder to cut individual DNA bands out of the gel for purifying, the present inventive method permits the user to simply pick DNA, together with agarose, out of the gel using a transfer pipette and soak the DNA in ddH₂O (about 50 µL) without purification. The DNA eluted from the agarose is sufficient for about at least 50 second-round polymerase chain reactions. Small amount of second PCR products can be saved when diluted in a large volume of buffer for a lifetime supply. A follow-up sequence identity check can usually confirm a product and remove any concerns about nonspecific PCR products or related sequences having similar size of the gene-specific products mixed together in the final products.

[0043] As mentioned before, two strategies are to be applied for amplification of the 3'UTR. The first strategy employs GSP with T7/T3 promoters for the first PCR, then use T7/T3 for the second PCR. An advantage of the first strategy is that it is able to simplify the procedures for a second round of PCR and subsequent sequencing verification of the final PCR products, because only a single pair of universal primers is required. Another advantage is having T7 and T3 promoters at both ends. Researchers will be able to generate RNA in either a sense or anti-sense direction, which ever and whenever necessary. The second strategy employs the same GSPs for both first and

second rounds of PCR. This approach has several advantages. It simplifies primer design, cuts the cost, and can avoid cross contamination problems. Additionally, the second strategy enables better verification of sequence, which provides a means for quality control of second-round PCR products since no PCR product will be generated if a mistake was made in mixing templates with primer pairs. No such control, however, will be associated with the first strategy because the universal primers can amplify any sequences from the first round of PCR.

Experiments

[0044] Experimental studies were conducted for 117 genes using the present method for amplifying expressed sequences from human genomic DNA. First, the relative size-distribution of the 3'UTR was ascertained according to the steps described above. The sequences for 117 putative tox genes were retrieved from the UniGene database and their respective 3'UTR were defined to determine how many genes have a 3'UTR length sufficient for PCR amplification. As shown in Figure 4, the 3'UTR for 29 genes are shorter than 200 bp (~24%), for 27 genes are between 200 to 400 bp (23%), and for 60 genes are over 400 bp (51%). Although the method can work with sequences of considerably less than 200 bp, such as short as 75-100 bp, a practical, minimal length required for PCR is about 200 bp. About 74% genes can be potentially amplified. Considering the constraints on sequence contents for primer design, 97 genes, each having a 3'UTR over 400 bp, were selected for PCR amplifications. [0045] Overall, two rounds of PCR were necessary to obtain sufficient DNA for array printing. The first round of PCR was carried out in a 10 μ L reaction volume under following conditions. Reagents: 1X buffer containing 1.5 mM MgCl₂ (PE Biosystems), 0.2 mM dNTP (GIBCO BRL), 0.4 µM of each primer, 100 ng human placenta genomic DNA, and 0.5 units of Taq polymerase (Roche Molecular Biochemicals). PCR cycles: one cycle of 95°C for 1 minute, 25 cycles of 94°C for 30 sec., 60°C for 30 sec., and 72°C for 45 sec., and one cycle of 72°C for 5 minutes. Gel electrophoresis was used to size-differentiate the PCR product on a 1.5% agarose gel. A transfer pipette picks up the DNA band with the expected size as defined by primer design software together with the slice of the gel on which the DNA rested, and placed the DNA in 50 μL water

to soak. One microliter of the DNA eluted out of the gel slice was used as templates

for second round of PCR using either T7/T3 primers or GSPs in 50 μ L reaction (8 reactions per gene) under the same condition described above. The PCR products (in total volume of 600 –1 for each gene) were cleaned using QIAquick PCR Purification kit (Qiagen), and eluted in a final volume of 100 μ L. One microliter of each product was loaded on a 1.5% agarose gel for verifying sizes and estimating concentrations. A randomly selected set of DNA samples was measured for OD₂₆₀ to set a standard for the adjustment of the DNA concentration for all PCR products.

[0046] The results from the first round of PCR amplification are shown in Figure 5A. Twelve genes were selected as proof of concept examples from the original 97 genes. The PCR products for the 12 genes that were amplified using Type I primers produced distinct, unique bands, each with the expected size. PCR, although a good tool, is still not sufficiently specific, nor perfect in amplifying correct sequences. The faint smear present in each lane of the gel represented nonspecific PCR products. Sizedifferentiation by gel electrophoresis, for instance, removes extraneous strands of a wrong sequence length. The wide DNA band observed near the loading well was from input genomic DNA. To remove nonspecific PCR products, a gel slice containing the DNA band of interest with correct length was removed and transferred to a tube containing about 50 µL of ddH₂O. The DNA eluted from the gel slice was then used for a second round of PCR. After electrophoresis column purification, 1 µL of each PCR product was again loaded on a gel for electrophoresis. Figure 5B shows the results from a second round of PCR using another 24 genes, also selected as examples of the original 97 genes, amplified using Type II primers. As seen in Figure 5B, all PCR products for the 24 genes gave a distinct single band, without visible background. All 12 genes amplified using the Type I primers, shown in Figure 5A, also gave the same results (data not shown). Generally, it was observed that once the first round of PCR amplification was done successfully, the second round of PCR would always work well, regardless the variations of the yield from gene to gene during the first round of PCR. In this particular experiment, over 90 percent of PCR products contained the correct sequence. In the field of microarray fabrication, an overall correct result of as high as over 90% is generally regarded as an excellent success rate for generating printable nucleic acid materials - especially in view of the difficulty of amplifying the kinds of genes selected herein.

[0047] Table 1 summarizes the results observed for both PCR products and sequencing. As recorded in Table 1, upper panel, a total of 97 genes were tried for PCR amplification. In the first round, the PCR products for 95 genes (95%) exhibited a distinct single band with their respective, expected size, and two genes ($\sim 2\%$) – BRAC2 (>900 bp) and CASP2 (>1200 bp) – had a single product longer than the cDNA sequence. The PCR products for three genes (~3%) – CASP13, COX11 and USP6 - had multiple bands from which no specific product could be identified. All PCR products were sequenced through the service provided by SeqWright Inc. Samples were prepared following manufacturer's instructions. Briefly, individual PCR products were diluted in ddH₂O to a final concentration of 50 ng/µL, and sequencing primers to 3.2 µM. The PCR products with either the correct size or wrong size for 94 genes were sequenced using a primer from sense direction. The results were summarized in the lower panel of Table 1. Briefly, the PCR products for 85 genes contain the correct sequences (90%); the sequences for 7 genes were not readable due to the presence of mixed sequences; and there were no signal for 2 genes probably due to sequencing system error (2%).

Table 1. Summary of Results Observed for PCR Products and Sequencing

	Gene M	Numbers (PCR	3)
Total	With expected size	With wrong size	No specific product
97	92 (95%)	2 (2%)	3 (3%)
	Gene Nun	nbers (Sequenc	cing)
Total	With correct sequence	Not readable	No signal
94	85 (90%)	7 (8%)	2 (2%)

The top panel shows the results of the first PCR; the bottom panel shows the results of sequencing. The percentile within parenthesis is calculated as follows: the number of genes within each category divided by the total number of genes shown in the first column.

[0048] Although the present invention has been described in detail, persons skilled in the art will understand that the invention is not limited to the embodiments specifically disclosed, and that various modification and variations can be made without departing from the spirit and scope of the invention. Therefore, unless changes otherwise depart from the scope of the invention as defined by the following claims, they should be construed as included herein.

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NM_002858 NM_012089 AF027302 NM_012089 AF027302 NM_001638 NM_001634 NM_001154 NM_001154 NM_001154 NM_001154 NM_001154 NM_001154 NM_00124 NM_001224 NM_001226 NM_001226 NM_001230 NM_001230 NM_001230 NM_001752 NM_001752 NM_001752 NM_001753 NM_001761 NM_004354 NM_004354 NM_004354 NM_004354 NM_004356 NM_004356 NM_004356 NM_004356 NM_004356 NM_004356 NM_004356 NM_004356 NM_004356	Symbol	Accesion No.	Sense primer	Antisense primer	Expected size, bp
NM_002858 NM_012089 AF027302 NM_001101 NM_001101 NM_001154 NM_001154 NM_001154 NM_001154 NM_001156 NM_000633 NM_000639 NM_001224 NM_001224 NM_001226 NM_001226 NM_001226 NM_001230 NM_001230 NM_001230 NM_001782 NM_001782 NM_001781 NM_004354 NM_004354 NM_004354 NM_004354 NM_004354 NM_004356 NM_004356 NM_004356 NM_004356 NM_004356 NM_004356	AATK	NM_004920	AATKs: T7-cttcactgactcagctagac*	AATKa: T3-accagcgttctaagcctcaa*	516
NM_001001 NM_001101 NM_001154 NM_001154 NM_001154 NM_001154 NM_0010574 NM_001059 NM_001224 NM_001224 NM_001226 NM_001227 NM_001230 NM_001237 NM_001731 NM_001732 NM_001732 NM_001732 NM_001731	ABCD3	NM_002858	ABCD3s: T7-tgactccaggaaaagccatt	ABCD3a: T3-tcgcttaggatcgtttgaca	537
AF027302 NM_001101 NM_001163 NM_001154 NM_001154 NM_001154 NM_001154 NM_001074 NM_001074 NM_000633 NM_000633 NM_001226 NM_001226 NM_001226 NM_001230 NM_001230 NM_001230 NM_001230 NM_001330 NM_001330 NM_001330 NM_001330 NM_001330 NM_001330 NM_001361 NM_001761 NM_001761 NM_001761 NM_001761 NM_001761 NM_001761 NM_001761 NM_001761 NM_0017661 NM_0017661 NM_0017661 NM_0017661 NM_0017661 NM_0017661 NM_0017661 NM_0017661 NM_0017661	ABCB10	NM_012089	ABCB10s: gcatggcacctcattttctt	ABCB10a: T3-agcagttcatgccttgcttc	484
NM_001101 NM_000668 NM_001635 NM_001154 NM_001154 NM_001154 NM_0010154 NM_001230 NM_001226 NM_001226 NM_001226 NM_001230 NM_001230 NM_001230 NM_001330 NM_001361 NM_001761 NM_001761 NM_001761 NM_001768 NM_001768 NM_001768 NM_001768 NM_001768 NM_001768 NM_001768 NM_001766 NM_001766 NM_001768 NM_001766 NM_001766	ABCFI	AF027302	ABCF1s, ateccactetgattgeatee	ABCF1a: gttcagcagtctttcc	408
NM_000668 NM_00154 NM_00154 NM_00154 NM_001674 NM_001674 NM_0010674 NM_0010674 NM_00126 NM_001226 NM_001226 NM_001226 NM_001230 NM_001230 NM_001231 NM_001231 NM_00124 NM_001252 NM_001254 NM_001261 NM_001261 NM_001761 NM_001761 NM_004354 NM_004354 NM_0043661 NM_001790 NM_001790 NM_001790	ACTB	NM_001101	ACTBs: T7-tgcgttacaccetttcttga	ACTBa: T3-gggagaccaaaagccttcat	541
NM_00154 NM_001154 NM_001168 NM_001168 NM_001674 AF021792 NM_0010574 NM_001059 NM_00124 NM_001226 NM_001226 NM_001230 NM_001230 NM_001231 NM_001231 NM_001231 NM_00124 NM_001251 NM_001251 NM_001251 NM_001761 NM_004354 NM_004354 NM_004354 NM_0040561 NM_0040561	ADH2	899000 WN	ADH2s: T7-gggccattgtgattgaagtc	ADH2a: T3-cattcacagcatttgccatc	559
NM_001154 NM_003734 NM_001168 NM_001674 AF021792 NM_0010633 NM_001026 NM_001224 NM_001226 NM_001226 NM_001227 NM_001230 NM_001752 NM_003723 NM_003723 NM_003724 NM_004354 NM_004354 NM_004354 NM_004354 NM_004661 NM_0040661 NM_0040661	AMPH	NM_001635	AMPHs: T7-ccctgcagaagatgtgatga	AMPHa: T3-tagectacetecagecacag	540
NM_003734 NM_001168 NM_001168 NM_001674 AF021792 NM_000633 NM_001224 NM_001224 NM_001226 NM_001226 NM_001226 NM_001226 NM_001727 NM_001730 NM_004354 NM_004354 NM_004354 NM_004354 NM_004661 NM_004661 NM_0047065	ANXA5	NM_001154	ANXA5s: T7-gcatttgtatgccagtgctt	ANXA5a: T3-ttcaggggacagaaatgtt	441
NM_001168	AOC3	NM_003734	AOC3s: T7-ccagagtagggttgccagtc	AOC3a: T3-attatcattgcaccccaaa	540
NM_001674 AF021792 NM_000633 NM_000196 NM_001059 NM_001224 NM_001224 NM_001226 NM_001227 NM_001227 NM_001230 NM_001752 NM_001730 NM_001751 NM_004761 NM_004661 NM_004661 NM_0047065	API4	NM_001168	API4s: T7-caggtgcctgttgaatctga	API4a: T3-aaggttgggctgacagacac	539
AF021792 NM_000633 NM_01196 NM_00196 NM_001324 NM_001224 NM_001226 NM_001227 NM_001230 NM_001527 NM_001527 NM_001530 NM_001752 NM_004354 NM_004354 NM_004354 NM_004661 NM_004661 NM_004661 NM_004661 NM_004661 NM_004661	ATF3	NM_001674	ATF3s: T7-ccagggttgtgctttctagc	ATF3a: T3-ctggtaccaccagctccact	527
NM_000633 NM_001196 NM_00059 NM_000344 NM_001224 NM_001226 NM_001226 NM_001227 NM_001230 NM_001752 NM_001752 NM_001761 NM_004354 NM_004354 NM_004661 NM_004661 NM_0047065	BAD	AF021792	BADs: T7-agtgacettegetecacate	BADa: T3-cagacgcgggctttattaac	417
NM_000196 NM_00059 NM_0004344 NM_001224 NM_001226 NM_001230 NM_001230 NM_001752 NM_001752 NM_001752 NM_001761 NM_004364 NM_004364 NM_004661 NM_004661 NM_004661	BCL2	NM_000633	BCL2As: T7-tggtgggaggaaaaggttg	BCL2Aa: T3-tctgagctccatcagcttcc	538
NM_000059 NM_004343 NM_001224 NM_001226 NM_001227 NM_001230 NM_001230 NM_001330 NM_001552 NM_005190 M74093 NM_004354 NM_004354 NM_004354 NM_0043661 NM_004661 NM_0017665	BID	961100_MN	BIDs: gaacggacagttccagaag	BIDa: tggaaataaaggcaccgtgt	293
NM_004343 NM_001224 NM_001326 NM_001227 NM_001230 C,NM_001330 NM_001752 NM_001752 NM_004354 NM_004354 NM_004354 C,NM_004354 NM_004354 C,NM_004354 NM_004361 NM_004361 NM_004661 CD	BRCA2	NM_000059	BRCA2s: T7-catttgcaaaggcgacaata	BRCA2a: T3-ctcaagtttgagtttggatgac	533
NM_001224 NM_004346 NM_001226 NM_001227 NM_001230 NM_001752 NM_001752 NM_0045190 NM_004354 NM_004354 NM_004354 NM_004661 NM_004661 NM_004665	CALR	NM_004343	CALRs: gogccaaataatgtctctgtg	CALRa: agaaagggagggtgaaatg	406
NM_004346 NM_001226 NM_001237 NM_001230 NM_001752 NM_001752 NM_001751 NM_001761 NM_004354 NM_004354 NM_004661 NM_001768 NM_004661 NM_007065	CASP2	NM_001224	CASP2s: gactgatcgtggggttgac	CASP2a: agaacagaaaccgtgcatcc	482
NM_001226 NM_001227 NM_001723 NM_001752 NM_001761 NM_001788 NM_001790 NM_001790 NM_001790 NM_001790 NM_001790 NM_001790 NM_001790	CASP3	NM_004346	CASP3s: catggtcaaaggctcaaacc	CASP3a: catglctctgctcaggctca	528
NM_001227 NM_001230 NM_003723 NM_005190 M74093 NM_001761 NM_001788 NM_004661 NM_001790 NM_007065	CASP6	NM_001226	CASP6s: ccaggcgfggftactcaca	CASP6a: ccatggccaacatgaacttt	427
NM_001230 NM_003723 NM_005190 NM_005190 NM_001761 NM_004354 NM_004388 NM_004661 NM_004661 NM_007065	CASP7	NM_001227	CASP7s: tccactgcaattggtggtaa	CASP7a: tggctttgttcttgtcatgg	200
NM_003723 NM_001752 NM_005190 M74093 NM_001761 NM_004354 NM_001788 NM_004661 NM_001790 NM_007065	CASP10	NM_001230	CASP10s: caggcaaagcttgaatcagg	CASP10a. cacctggctgaagtcaaatc	509
NM_001752 C NM_005190 C M74093 C NM_001761 C NM_004354 C NM_004661 C NM_001790 C NM_001790 C NM_007065 C C	CASP13	NM_003723	CASP13s: cagggtgaaaggagatggtg	CASP13a: aagtggtacatctccttagtc	497
NM_005190 M74093 NM_001761 NM_004354 NM_004661 NM_001790 NM_001790	CAT	NM_001752	CATs: taaccegeteateactggat	CATa attaagccatgacggtgctc	445
M74093 NM_001761 NM_004354 NM_001788 NM_001790 NM_001790 CI	CCNC	NM_005190	CCNCs: aaacattccgaagaattcca	CCNCa: ggtccctcaafgaccaaaga	376
NM_001761 NM_004354 NM_001788 NM_004661 NM_001790 NM_007065	CCNEI	M74093	CCNE1s: ccatcettetecaceaaga	CCNE1a: ctatgggctctgcacaacg	403
NM_004354 NM_001788 NM_004661 NM_001790 CI	CCNF	NM_001761	CCNFs: getgecatecacttetgttt	CCNFa: ggtggccagaattcccttat	501
NM_001788 NM_004661 NM_001790 CI	CCNG2	NM_004354	CCNG2s: agccatcaaatggggtagtg	CCNG2a: cttggggcaataggaatgaa	501
NM_004661 NM_001790 CDC NM_007065	CDC10	NM_001788	CDC10s: caaaggttccattcagtgcag	CDC10a: cttcaagaggccatgattcc	491
NM_001790 NM_007065	CDC23	NM_004661	CDC23s: gaccttgetettggatttge	CDC23a: acaggcctgaaactctccaa	505
NM_007065	CDC25C	NM_001790	CDC25Cs: ggctgctaacaagtcaccaaa	CDC25Ca: caacgetettgeatagece	324
	CDC37	NM_007065	CDC37s: ctgcttccagccctatgt	CDC37s: gacacagacagacgaaca	340
NM_001791	CDC42	NM_001791	CDC42s: gacaaatgcctgcacctac	CDC42a: caatccgtcctccctta	422

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CDKN2Aa: gccatttgctagcagtgtga 414	CHES1a: gccaatcttcaggcttatgg 501	CLGNa. tgaacaaggcatgtccttaaa	COX10a: ccagcacacccttcttccta 502	COX11a: attetttaggggccaggatc 480	COX15a: cagctctgcagcataatgga 496		CYP1B1a: ccatgctttgaattttgtgc 509	CYP3A3a: tgtcattgttagagccatcaaaa 320	CYP4A11a: tgtgacggtttagcatctgc 499	CYP4B1a: catctcagtgaagggggact 426	CYP4F2a: gtgtcgtgctaccttcgtca 492	CYP4F3a, tcaccatcccaggagaaaac 497	CYP7A1ar atgatcacaccegaagaacc 499	CYP7B1a: gggaaacattttcatccagtg 439	CYP8B1a: ttggagaaagctggcaaagtt 500	CYP19ar ccccaatcactgtagctgt 506	CYP24a: caaataatgccccagtgaatc 510	CYP51a: gaagcaggaacaactgagc 503	DAPK3a: attectetggetgcagagg 443	DHFRa: atgcaaccetttggttcaag 499	DNJ3a: gccaaacacaaagcttcagg 385	DPYDa: tgaagatgccatgaagagga 481	DTRa: cagctccaatgttccctgtt 493	EGFa: tgtgcaatcacacaagagg 461	EGR1a: catgtccctcacaattgcac 501	EPOa: gtcttcatggttccaccac 453	FADDa: ttgcaggacccataatcctc 506	G6PDa: tagcagagaggctgcctacg 455	G17a: cctgtttcttctccagcag	GAS11a: ctctgggcctaacctcactg 500	HIF1Aa: gcgacaaagtgcataaaatcaa	
CDKN2Aa: g	CHES1a: g	CLGNa. 1g	COX10a: c	COX11a: a	COX15a: ca	CPT2a:	CYPIBIa:	CYP3A3a: tgtc	CYP4A11a: t	CYP4B1a: ca	CYP4F2a: g	CYP4F3a, to:	CYP7A1a at	CYP7B1a: gg	CYP8B1a: ttg	CYP19a c	CYP24a: ca	CYP51a: gaa	DAPK3a: a	DHFRa: a	DNJ3a: go	DPYDa: tga	DTRa: 0	EGFa: tgt	EGR1a: c	EPOa: g	FADDa: tt.	G6PDa: tag	G17a: c	GAS11a: ct	HIF1Aa: gcga	HPRT1a: gggaactoctoacaagatte
CDKN2As: totgagaaacctcgggaaac	CHES1s: cctccagcttgtcagaaacc	CLGNs: agcatgccagacctgaactt	COX10s: gtgagcctcatgatctgctg	COX11s tcacgctgttgtcaggaatc	COX15s: tgacccafcgagatgaaat	CPT2s: getaccateaetteeteate	CYP1B1s: tggggacagaactcccatta	CYP3A3s: gcctgagaacaccagagacc	CYP4A11s: cctgtctgccatatcctgt	CYP4B1s: atgagaatggggtccagat	CYP4F2s: ccctaagaccctgttccaca	CYP4F3s: cccactaaaatgaccctga	CYP7A1s: ttgttcaccagtgcttgctt	CYP7B1s: ccctaaacatcctaagctcatct	CYP8B1s: ettetatececagacecae	CYP19s: ccaaacccacctgctagtgt	CYP24s: tgggatccaaggcattctac	CYP51s: actcatcgctcttgccaaat	DAPK3s: gggctgcttctctacacagc	DHFRs: gggaacagtgaatgccaaac	DNJ3s: ctgcaaacaaattgcacagg	DPYDs: cccttcgctgaaattgctta	DTRs: cctttgccacaaagctagga	EGFs: caaattgggacaacagtgctt	EGR1s: ccttgctcccttcaatgcta	EPOs: ctcctcaccaacattgctt	FADDs: tgcgggagtagttggaaagt	G6PDs: ttgacctcagctgcacattc	G17s: ctaccetgetaggetetgg	GAS11s: gaatggacagctttgcaggt	HIF1As: gtggtagccacaattgcaca	HPRTIS: aetteteteecateteett
NM_000077	NM_005197	NM_004362	NM_001303	NM_004375	NM_004376	NM_000098	NM_000104	NM_000776	NM_000778	NM_000779	NM_001082	968000 ⁻ MN	NM_000780	NM_004820	AF090320	NM_000103	NM_000782	NM_000/86	NM_001348	NM_000791	NM_004222	NM_000110	NM_001945	NM_001963	NM_001964	NM_000799	NM_003824	NM_000402	NM_006841	NM_001481	NM_001530	NM_000194
CDKNZA	CHESI	CLGN	COX10	COXII	COXIS	CPT2	CYPIBI	CYP3A3	CYP4A11	CYP4B1	CYP4F2	CYP4F3	CYP7A1	CYP7B1	CYP&B1	CYP19	CYP24	CYPSI	DAPK3	DHF.K	DNJ3	DPYD	DIR	EGF	EGRI	EPO	FADD	GePD	617	GASH	HIFIA	IIIIII

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IGF2R	NM_000876	IGF2RS: attegaggaageettecte	april 7	
ITGA5	NM 002205	TTGA Sc. moonstatt	ICF2Ra: atctttgggcaggttgttg	206
LPL	NM 000237	I DI O COLOR	ITGA5a: ggaaattcctggcttctcct	493
MADD	NM 003682	Lr LS. tatagctgggacccgactg	LPLa: gccacaatgacctttccaat	506
MADH2	100500 MN	MADELS acceptualgreectorg	MADDa: cgaccactccatctgat	507
MAOB	NM 000898	MADH28: caatcaagtcccatggaaaag	MADI12a: atcaagaagcagcacac	397
MAP3K8	NM 005204	MADING: ttccaagttattgccctcaa	MAOBa: agacacaccacaaaacag	504
MMP14	NM 004995	MANDIA.	MAP3K8a: tcactagtggccgtctgtca	501
NAT2	NM 000015	MATT2	MMP14a: tcgtttgtgtgccttctctg	499
IGON	NM 006092	NATAS: cellgigatgateacecaacte	NAT2a: agcatgaatcactctgcttcc	243
NR112/SXR	AJ009937	ND 172:	NOD1a: ccatgccctatttctttgga	502
PDCDI	NM 005018	DDCD15: cacatacccacguigueg	NR112a: tgcccttgctcctacagact	206
RAD9	NM 004584	DATA.	PDCD1a: ggaccgtaggatgtcctct	200
RBI	NM 000321	D.D.J.:	RAD9a: agcgccaaagagtatcagga	495
REQ	NM 006268	N.D. IS: Igaggaccttgg	RB1a: gtgaatgggcagtcaatcaa	486
SLC15A1	NM 005073	CT C15 A 1	RAQa: tcaactccaaagcgacagtg	496
SLC20A2	NM 006749	SLC13A1S: Uctaagcagcagga	SLC15A1a: toattactcggccttcacct	411
SLC29A1	NM 004955	SLC-20A43. gcaaacagctaaagggatgg	SLC20A2a: ggttgcctgttctgaagctc	480
SMAC	73355 NN	SLC29A1S: ggtgatcctgagtggtctgg	SLC29A1a: aaggcacctggtttctgtca	306
TNFRSF6/FAS	NM 000043	SMACS: tgtctgtgcaccgagaagag	SMACa: cctgttgagagcaccaggta	505
TNFSF6	059000 MN	INFRNF6S: tagagetttgccacctctcc	TNFRSF6a: ggtggttccaggtatctgct	506
TP53	NM 000546	1 NFSF68: tgttacaggcaccgagaatg	TNFSF6a: gttagttfcaccgatggctc	488
UCH37	NM 015984	11555. CCugcilgcanaggigt	TP53a: tacctaaccagctgcccaac	502
UGTIAI	NM 000463	UCH3/S: gettetgeacatatttteatgg	UCH37a: tcactggaaattatacttitgtccttt	510
USP5	NM 003481	UGITAIS: taatcagccccagagtgctt	UGT1A1a: acaccaccaccaatttcat	480
9dSI1	MM O04505	USP5S: ctfaccaafgagggcaggg	USP5a: ggcatttccagagaagaca	503
8dS11	COCEDO TAINI	USP6s: taatagcagcccacggactt	USP6a: ggcagagtcggtgtcaattt	505
0.00	MW_003134	USP8s: aggacagtgggagctgtgtt	USP8a: atасарсссаяарс	777
USF11	NM_004651	USP11s: cetetetgeaatetegette	[[Sp11a; aggaanateacteath	1/4
	NM_005151	USP14s: cacccaagattcagcagtca	110D140: etattee	/55
USP15	AF106069	1100.1	Cor 14a. gicilicagecaagetecaae	490

* Type I primers with T7 or T3 promoter at the 5'end.